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### (54) v-TRIAZOLYL-[4,5-d]-PYRIMIDINES AND THEIR USE AS OPTICAL BRIGHTENING AGENTS FOR ORGANIC MATERIALS

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(71) We, CIBA-GHIGY AG, a Swiss body corporate, of Basle, Switzerland, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

The present invention relates to v-triazolyl-[4,5-d]-pyrmidines, their preparation and their use for the optical brightening of organic,

especially polymeric, materials.

The v-triazolyl[4,5-d]-pyrimidines of the invention have the formula

in which Q and Q<sub>1</sub> independently of one another are a secondary or tertiary amino radical; X is hydrogen, alkyl having 1 to 6 carbon atoms, alkoxy having 1 to 6 carbon atoms, allyloxy, hydroxyalkoxy having 2 to 4 carbon atoms, alkoxyalkoxy having a total of 3 to 8 carbon atoms, benzyloxy, phenethyloxy, halogen, phenoxy, phenoxyalkoxy having 1 to 3 carbon atoms in the alkoxy part, sulpho, cyano, —COOY (in which Y is hydrogen, a salt-forming cation or alkyl having 1 to 18 carbon atoms) or

-SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or -CONY<sub>1</sub>Y<sub>2</sub>

(in which  $Y_1$  and  $Y_2$  independently of one

another are hydrogen or alkyl having 1 to 4 carbon atoms); X<sub>1</sub> is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms, halogen, sulpho, cyano,—COOY (in which Y is hydrogen, a saltonning cation or alkyl having 1 to 18 carbon atoms) or

-SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or -CONY<sub>1</sub>Y<sub>2</sub>

(in which Y<sub>1</sub> and Y<sub>2</sub> independently of one another are hydrogen or alkyl having 1 to 4 carbon atoms) or X and X<sub>1</sub> together form a methylenedioxy, ethylenedioxy or methyleneoxymethyleneoxy radical; and X<sub>2</sub> is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms or halogen.

Secondary and tertiary amino radicals Q and Q<sub>1</sub> are to be understood as meaning radicals which are of the formulae —NR°R<sub>1</sub>° and —NR<sub>2</sub>°R<sub>3</sub>° respectively, in which R° and R<sub>1</sub>° and, respectively, R<sub>2</sub>° and R<sub>2</sub>° independently of one another are unsubstituted alkyl having 1 to 12, and preferably 1 to 4, carbon atoms, hydroxyalkyl having 2 to 12, and preferably 2 to 6, carbon atoms, alkyl having 2 to 4 carbon atoms which is substituted by alkoxy having 1 to 8 carbon atoms or dialkylamino having 2 to 4 carbon atoms per alkyl part, or together with the nitrogen atom are a 5-membered or 6-membered saturated heterocyclic structure which can contain other hetero-atoms, and R° and R<sub>2</sub>° are also hydrogen.

Possible heterocyclic structures are, for example, pyrrolidine, piperidine, piperazine and morpholine, which can be unsubstituted

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or substituted by alkyl having 1 to 4 carbon atoms or halogen. Piperazine rings can be substituted in the 4-position by alkyl or hydroxyalkyl having 1 to 4 carbon atoms or can also be quaternised.

Sulpho is to be understood as meaning, in each case, the radical -SO,M, in which M is hydrogen or a salt-forming cation. Possible cations are, in general, those of alkaline earth 10 metals, for example of calcium, barium or magnesium, and also especially of alkali metals, for example of sodium or potassium.

Important compounds within the scope of the compounds of the formula (1) are those of the formulae

in which R and R<sub>2</sub> independently of one another are hydrogen, alkyl having 1 to 12 carbon atoms, hydroxyalkyl having 2 to 6 carbon atoms or dialkylaminoalkyl having 2 to 4 carbon atoms per alkyl part; R1 and R, independently of one another are alkyl having 1 to 12 carbon atoms or hydroxyalkyl having 2 to 6 carbon atoms; or R<sub>1</sub> together with R, or R, together with R<sub>2</sub>, form with the nitrogen atom to which they are attached 5-membered or 6-membered saturated heterocyclic structure which can contain one or more further hetero-atoms; X is hydrogen, alkyl having 1 to 6 carbon atoms, alkoxy having 1 to 6 carbon atoms, allyloxy, hydroxyalkoxy having 2 to 4 carbon atoms, alkoxy-alkoxy having a total of 3 to 8 carbon atoms, benzyloxy, phenethoxy, halogen, phenoxy, phenoxyalkoxy having 1 to 3 carbon atoms in the alkoxy part, sulpho, cyano, —COOY, or —SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or CONY<sub>1</sub>Y<sub>2</sub> and X<sub>1</sub>' is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms, halogen, sulpho, cyano, —COOY, or X and X, form a methylenedioxy, ethylenedioxy methyleneoxymethyleneoxy radical, and

in which R' and R2' independently of one another are hydrogen, alkyl having 1 to 4 carbon atoms or hydroxyalkyl having 2 to 6 carbon atoms; R,' and R,' independently of one another are alkyl having 1 to 4 carbon atoms or hydroxyalkyl having 2 to 6 carbon

atoms, or R1' together with R', or R1' together with R, rogether with the nitrogen atom to which they are attached, form a 5-membered or 6-membered saturated heterocyclic structure which can contain a further nitroger atom or an oxygen atom; X' is hydrogen, alky having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms, allyloxy, hydroxyalkoxy having 2 to 4 carbon atoms, alkoxy-alkoxy having a total of 3 to 8 carbon atoms, benzyloxy, phenethoxy, chlorine, phenoxy, phenoxy-propoxy, sulpho, cyano, —COOY, and X<sub>1</sub>" is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms or chlorine, or  $X_1''$  and X' together form a methylenedioxy, ethylenedioxy or methyleneoxymethyleneoxy

Compounds of particular interest are vtriazolyl [4,5-d]-pyrimidines of the formula

in which R" and R<sub>2</sub>" independently of one another are hydrogen, alkyl having 1 to 4 carbon atoms or hydroxyethyl; R<sub>1</sub>" and R<sub>2</sub>" independently of one another are alkyl having 1 to 4 carbon atoms or hydroxyethyl; or R<sub>1</sub>" together with R", or R<sub>3</sub>" together with R<sub>4</sub>", and the nitrogen atom to which they are attached are pyrrolidino, piperidino, morpholino, N'-hydroxyethylpiperazino or N'-hydroxyethylpiperazino quaternised by alkyl having 1 to 4 carbon atoms, and X" is hydrogen, methyl, chlorine, alkoxy having 1 to 4 car-bon atoms, allyloxy, hydroxyalkoxy having 2 to 4 carbon atoms, alkoxyalkoxy having to 6 carbon atoms, antoyatovy naving 5 to 6 carbon atoms, benzyloxy, phenoxy, phenoxy, sulpho, cyano, —COOY (in which Y is hydrogen, a salt forming cation or alkyl having 1 to 4 carbon atoms), or —SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or —CONY<sub>1</sub>Y<sub>2</sub>, X<sub>1</sub>" is hydrogen, a salt of the salt of gen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms or chlorine, or X" and X," together form a 3,4-methylenedioxy or 3,4-ethylenedioxy or 3,4-methyleneoxymethyleneoxy radical.

Compounds of particular importance are

v-triazolyl-[4,5-d]-pyrimidines of the formula

in which R" is alkyl having 1 to 4 carbon atoms or two adjacent R" together with the nitrogen atom to which they are attached are piperidino or morpholino; X" is hydrogen, 100

methyl, allyloxy, hydroxyethoxy, alkoxyalkoxy baving a total of 3 to 6 carbon atoms, benzyloxy, alkoxy having 1 to 4 carbon atoms, phenoxy, phenoxypropoxy or sulpho, and X''' is hydrogen, alkyl having 1 to 4 carbon atoms or methoxy, or K," and K" together form a methylenedioxy, 3,4-ethylenedioxy or 3,4-methyleneoxymethyleneoxy, radical.

The v-triazolyl[4,5-d] pyrimidines of the formula (1) can be prepared by processes which are known per se. They can be obtained, for example, by a process which com-prises coupling an amine of the formula

in which Q and Q, are as defined above, with a diazonium salt of the formula

in which Z is an anion and X, X, and X, are as defined above, and oxidatively cyclising the resulting o-aminoazo compound of the formula

(8) 
$$Q_1 \xrightarrow{N}_{M} N = N \xrightarrow{X}_{M_2} X_1$$

in which Q, Q1 and X to X2 are as defined above, at a temperature of from 70 to 100° C. The oxidative cyclisation can be effected in a manner known per se, e.g. from U.S. Patent Specification 2,543,333.

Possible anions Z are, for example, a halogen ion, such as a chloride or bromide ion, a

sulphate ion or a fluoborate ion.

Some of the amines in which X is a radical bonded via an oxygen atom to the benzene ring, which are required for the preparation of the diazonium salts of the formula (7), are known and some are novel. The novel amines can be prepared by known methods, for example by reacting p-nitro-chlorobenzene with an alcohol and subsequently reducing the nitro group to the amino group. Suitable amines of this type are, for example, 1 - methoxy - 2 - (4 - aminophenoxy) - ethane, 1 - isopropoxy - 2 - (4aminophenoxy) - ethane, 1 - n - butoxy - 2-(4 - aminophenoxy) - ethane, 1 - methoxy-2 - (4 - aminophenoxy) - isopropane, 1-hydroxy - 2 - (4 - aminophenoxy) - ethane or 2 - methoxy - 4 - (4 - aminophenoxy)sec - butane.

Oxidising agents which can be used to effect the oxidative cyclisation are, for example, chromic acid, alkali metal bichromates, hydrogen peroxide, lead tetrascetate, potassium ferricyanide, ferric chloride and copper-II sulphate. In acid solvents (for example aqueous acetic acid), alkali metal bichromates, hydrogen peroxide or lead tetraacetate are preferably used and in basic solvents (for example a pyridine/water mixture) potassium ferricyanide is preferably used. The oxidative cyclisation is preferably effected with copper-II sulphate in a pyridine/water mixture. The oxidation with copper-II salts, such as copper-II sulphate or copper-II chloride, can advantage-ously also be carried out in methanol or methanol/water mixtures in the presence of ammonium salts or amine salts, such as monoalkanolamines or dialkanolamines. The coupling of an amine of the formula (6) with a diazonium salt of the formula (7) is usually carried out at a temperature of -10 to 20° C, preferably of 0 to 10° C. The oxidative cyclisation is effected at a temperature of from 70 to 100° C, preferably 90 to 100° C.

In the dissolved or finely divided state, the compounds of the invention have a more or less pronounced fluorescence. They can be used for the optical brightening of synthetic, semi-synthetic or natural organic materials or of substances which contain such organic materials.

The following groups of organic materials,

where optical brightening thereof is relevant, may be mentioned as examples of the above without the survey given below being intended to express any restriction thereto:

#### Synthetic organic high-molecular materials:

polymerisation products based on organic compounds which contain at least one polypolymerisable carbon-carbon double bond, i.e. the homopolymers or copolymers thereof and also the after-treatment products thereof, for example, crosslinking products, grafting products or degradation products, polymer blends or products obtained by modifying reactive groups, for example polymers based on  $\alpha_s\beta$ unsaturated carboxylic acids or derivatives of such carboxylic acids, and especially on acrylic compounds (for example acrylates, acrylic acid, acrylonitrile, acrylamides and the derivatives thereof or the methacrylic analogues thereof) and on olefine hydrocarbons (for example ethylene, propylene, styrenes or dienes, and also so-called ABS polymers), and polymers based on vinyl and vinylidene compounds (for example vinyl chloride, vinyl alcohol and vinylidene chloride).

b) polymerisation products which are obtainable by ring opening, for example polyamides of the polycaprolactam type, and also 110

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polymers which are obtainable by polyaddition or by polycondensation, such as polyethers or

polyacetals,

c) polycondensation products or precondensates based on bifunctional or polyfunctional compounds having groups capable of undergoing a condensation reaction, the homocondensation and cocondensation products thereof and also after-treatment products, for example polyesters, especially saturated (for example ethylene glycol terephthalic acid polyesters) or unsaturated (for example maleic acid/dialcohol polycondensates and also their crosslinking products with copolymerisable vinyl monomers), non-branched and branched (also those based on polyhydric alcohols, for example alkyd resins) polyesters, polyamides (for example hexamethylenediamine adipate), maleate resins, melamine resins, their precondensates and analogues, polycarbonates and silicones, and

d) polyaddition products such as polyurethanes (crosslinked and non-crosslinked) and

epoxide resins.

II. Semi-synthetic organic materials, for example cellulose esters of various degrees of esterification (80-called 2½-acetate and tri-acetate) or cellulose ethers, regenerated cellulose (viscose or cuprammonium cellulose), or their after-treatment products, and casein plastics.

III. Natural organic materials of animal or vegetable origin, for example those based on cellulose or proteins, such as cotton, wool, linen, silk, natural lacquer resins, starch and

casein.

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The organic materials to be optically brightened can be in various different states of processing (raw materials, semi-finished goods or finished goods). On the other hand, they can be in the form of structures of very diverse shapes, i.e., for example, in the form of predominantly three-dimensional bodies, such as slabs, profiles, injection mouldings, various machined articles, chips, granules or foams, and also in the form of predominantly two dimensional bodies, such as films, sheets, lacquers, coverings, impregnations and coatings, or in the form of predominantly one-dimensional bodies such as filaments, fibres, flocks and wires. The said materials can, on the other hand, also be in an unshaped state, in very diverse homogeneous or inhomogeneous forms of division, for example in the form of powders, solutions, emulsions, dispersions, latices, pastes or waxes.

Fibre materials can, for example, be in the form of continuous filaments (stretched or unstretched), staple fibres, flocks, hanks, textile filaments, yarns, threads, fibre fleeces, felts, waddings or flocked structures or in the form of woven textile fabrics or textile laminates and knitted fabrics, and also in the form of papers, cardboards or paper pulps.

The compounds to be used according to the invention are of importance, inter alia, for the treatment of organic textile materials, especially woven textile fabrics. Where fibres, which can be in the form of staple fibres or continuous filaments or in the form of hanks, woven fabrics, knitted fabrics, nonwovens, flocked substrates or laminates; are to be optically brightened according to the invention, this is advantageously effected in an aqueous medium, in which the compounds in question are present in a finely divided form (suspensions, so-called microdispersions or possibly solutions). If desired, dispersing agents, stabilisers, wetting agents and further auxiliaries can be added during the treatment.

Depending on the type of brightener compound used, it can prove advantageous to carry out the treatment in a neutral or alkaline or acid bath. The treatment is usually carried out at temperatures of from 20 to 140°C, for example at the boiling point of the bath or near it (about 90° C). Solutions or emulsions in organic solvents can also be used for the finishing, according to the invention, of textile substrates, as is practised in the dveing trade in so-called solvent dyes (padthermofix application, or exhaustion dye processes in dyeing machines).

The optical brighteners according to the present invention can furthermore be added to, or incorporated in, the materials before or during their shaping. Thus they can, for example, be added to the compression moulding compositions or injection moulding compositions during the production of films, sheets (for example hot milling into polyvinyl

chloride) or mouldings.

Where fully synthetic or semi-synthetic organic materials are being shaped by spinning processes or via spinning compositions, the optical brighteners can be applied in accordance with the following processes:

addition to the starting substances example monomers) or intermediates (for example precondensates or prepolymers), i.e. before or during the polymerisation, polycondensation or polyaddition,

powdering onto polymer chips or granules for spinning compositions,

bath dyeing of polymer chips or granules 115 for spinning compositions, metered addition to spinning melts or

spinning solutions, and

application to the tow before stretching. The optical brighteners according to the 120 present invention can, for example, also be employed in the following use forms:

a) mixed with dyes (shading) or pigments (coloured pigments or especially, for example, white pigments), or as an additive to dye baths, printing pastes, discharge pastes or reserve pastes, and also for the after-treatment of dyeings, prints or discharge prints,
b) mixed with so-called "carriers",

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wetting agents, plasticisers, swelling agents, anti-oxidants, light stabilisers, heat stabilisers and chemical bleaching agents (chlorite bleach

or bleaching bath additives),

c) mixed with crosslinking agents or finishing agents (for example starch or synthetic finishes) and in combination with the most diverse textile finishing processes, especially synthetic resin finishes (for example crease-proof finishes such as "wash-and-wear", "permanent-press" or "no-iron"), and also flame-proof finishes, soft handle finishes, antisoiling finishes or antistatic finishes, or antimicrobial finishes,

d) incorporation of the optical brighteners into polymeric carriers (polymerisation, poly-condensation or poly-addition products), in a dissolved or dispersed form, for use, for example, in coating agents, impregnating agents or binders (solutions, dispersions and emulsions) for textiles, fleeces, paper and leather,

e) as additives to so-called "master batches",

25 f) as additives to the most diverse industrial products in order to render these more marketable (for example improving the appearance of soaps, washing agents and pig-

in combination with other optically

brightening substances,

h) in spinning bath formulations, i.e. as additives to spinning baths, such as are used for improving the slip for the further processing of synthetic fibres, or from a special bath before stretching the fibre,

i) as scintillators for various purposes of a photographic nature, for example for electrophotographic reproduction or supersensitisa-

j) depending on the substitution, as laser

dyes

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If the brightening process is combined with textile treatment methods or finishing methods, the combined treatment can in many cases advantageously be carried out with the aid of appropriate stable formulations which contain the optically brightening compounds in such concentration that the desired brightening effect is achieved.

In certain cases, the brighteners are made fully effective by an after-treatment. This can, for example, be a chemical treatment (for example acid treatment), a thermal treatment (for example heat) or a combined chemical/thermal treatment. Thus, example, the appropriate procedure to follow in optically brightening a range of fibre substrates, for example polyester fibres, with the brighteners according to the invention is to impregnate these fibres with the aqueous dispersions (or where appropriate also solutions) of the brighteners at temperatures below 75° C, for example at room temperature, and to subject them to a dry heat treatment at tempera-

tures above 100° C, it generally being advisable additionally to dry the fibre material beforehand at a moderately elevated temperature, for example at not less than 60° C and up to 130° C. The heat treatment in the dry state is then advantageously carried out at temperatures between 120 and 225° C, for example by heating in a drying chamber, by ironing within the specified temperature range or by treatment with dry, superheated steam. The drying and dry heat treatment can also be carried out in immediate succession or be combined in a single operation.

The amount of the optical brighteners to be used according to the invention, relative to the material to be optically brightened, can vary within wide limits. A distinct and durable effect is already achievable with very small amounts, in certain cases, for example, amounts of 0.0001 per cent by weight. Usually, at least 0.001 per cent by weight is advisable. Amounts of up to 0.8 per cent by weight and in some cases of up to 2 per cent by weight can also be employed. For most practical purposes, amounts between 0.0005 and 0.5 per cent by weight are of preferred interest.

For various reasons it is frequently appropriate to employ the brighteners not in the form of the compounds themselves, i.e. as the pure compounds, but as a mixture with very diverse auxiliaries and diluents, for example anhydrous sedium sulphate, sedium sulphate decahydrate, sedium chloride, sedium carbonate, alkali metal phosphates, such as sodium orthophosphate or potassium orthophosphate, 100 sodium pyrophosphate or potassium pyro-phosphate and sodium tripolyphosphates and potassium tripolyphosphates, or alkali metal ailicates.

The optical brighteners are also particularly 105 suitable for use as additives for wash liquors or industrial and domestic washing agents, to which they can be added in various ways. They are appropriately added to wash liquors in the form of their solutions in water or organic 110 solvents or in a finely divided form, as aqueous dispersions. They are advantageously added to domestic or industrial washing agents at any stage of the production process of the washing agent, for example to the so-called "alurry" before spray-drying, to the washing powder, or during the preparation of liquid washing agent combinations. They can be added either in the form of a solution or dispersion in water or other solvents or, without auxiliaries, as a dry brightener powder. For example, the brighteners can be mixed, kneaded or ground with the detergent sub-stances and, in this form, mixed into the finished washing powder. They can, however, be sprayed in a dissolved or predispersed form onto the finished washing agent.

Possible washing agents are the known mixtures of detergent substances, for example soap in the form of chips and powders, syn-

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thetics, soluble salts of sulphonic acid halfesters of higher fatty alcohols, arylsulphonic acids having higher and/or multiple alkyl substituents, sulphocarboxylic acid esters of medium to higher alcohols, fatty acid acylaminoalkyl- or acylaminoaryl-glycerolsulphonates, and phosphoric acid esters of fatty alcohols. Examples of so-called "builders" which can be used are alkali metal polyphosphates and polymetaphosphates, alkali metal pyrophosphates, alkali metal salts of carboxymethylcellulose and other "soil redeposition inhibitors", and also alkali metal silicates, alkali metal carbonates, alkali metal borates. alkali metal perborates, nitrilotriacetic acid, ethylenediaminetetraacetic acid and foam stabilisers, such as alkanolamides of higher fatty acids. The washing agents can also contain, for example: antistatic agents, superfatting skin protection agents, such as lanolin, enzymes, antimicrobial agents, perfumes and dyes.

The compounds of the invention have the particular advantage that they are also effective as optical brightening agents in the presence of active chlorine donors, for example hypochlorite, and can be used without significant loss of effect in wash liquors containing non-ionic washing agents, for example alkylphenol

polyglycol ethers.

The compounds according to the invention are added in amounts of 0.005—1% or more, relative to the weight of the liquid or pulverulent, finished washing agent. Wash liquors which contain the indicated amounts of the optical brighteners claimed impart a brilliant appearance in daylight when used to wash textiles of cellulose fibres, polyamide fibres, cellulose fibres with a high quality finish, polyester fibres and wool.

The washing treatment is carried out, for

example, as follows:

The cited textiles are treated for 1 to 30 minutes at 20 to 100° C in a wash liquor which contains 1 to 10 g/kg of a composite washing agent containing a builder and 0.05 to 1%, relative to the weight of the washing agent, of the brighteners claimed. The liquor ratio can be 1:3 to 1:50. After washing, rinsing and drying are carried out as usual. The wash liquor can contain 0.2 g/l of active chlorine (for example in the form of hypochlorite) or 0.1 to 2 g/l of sodium perborate, as a bleaching additive.

In the examples, percentages are always percentages by weight. Unless otherwise stated, the melting points and boiling points are un-

corrected.

Example 1.

5.9 g of p-toluidine are dissolved in 140 ml

of water and 16.5 ml of concentrated hydrochloric acid, with stirring, the solution is cooled to 0 to 5° C and a solution of 3.8 g of sodium nitrite in 10 ml of water is added in the course of 15 to 20 minutes at this temperature. The solution is then stirred for a further 30 minutes at 0 to 5° C and added dropwise, at 0 to 5° C, to a solution, which has been pre-cooled to 0° C, of 10.0 g of 4 - amino - 2,6 - his - (dimethylamino)-pyrimidine in 100 ml of pyridine, whereupon an exothermic reaction takes place and a yellow suspension forms. After all of the solution has run in, the resulting suspension is left to stand for 6 hours at 0 to 5° C and is then stirred for 12 hours at 20° C. The product formed is filtered off, washed with water and dried in vacuo at 70° C. This gives 16.3 g (99% of theory) of the yellow azo material having a melting point of 172 to 174° C.

15.0 g of the resulting compound are dissolved in 125 ml of pyridine at 60° C, with stirring, and a solution of 31.3 g of copper-II sulphate pentahydrate in 125 ml of water is added at this temperature. The mixture is then stirred under reflux for 4 hours and cooled to 20° C and the product is filtered off, washed well with water and dried in vacuo at 100° C. Recrystallisation from chlorobenzene while treating with bleaching earth gives 10.7 g (72% of theory) of the compound of the formula

in the form of pale yellow crystals. Melting point: 209 to 210° C.

The compounds of the formula

which are listed in Table I can be prepared in a manner similar to that described above 1 using the corresponding 4 - amino - 2,6diamino - pyrimidines and substituted anilines

TABLE

No.	R	R <sub>1</sub>	x	X <sub>1</sub>	Melting point uncorrected in °C
(101)	Сн,	СН,	4-OCH,	Н	209-210
(102)	Сн,	Сн,	4-C1	H	270—272
(103)	CH,	CH,	3-C1	Н	198-199
(104)	СН,	CH <sub>2</sub>	Н	н	199-201
(105)	сн,	CH,	3-CH,	н	163—164
(106)	CH,	CH,	2-CH,	н	150-151
(107)	Сн,	СН,	3-0CH <sub>3</sub>	н	177.5—179
(108)	CH,	CH,	2-OCH <sub>3</sub>	н	188-189.5
(109)	CH,	CH,	4-0(CH <sub>2</sub> ) <sub>3</sub> 0-	н	135—137
(110)	CH,	СН,	4-0	н	216—217
(111)	СН,	CH,	4-O(CH <sub>2</sub> ), CH <sub>3</sub>	н	173—174
(112)	CH <sub>3</sub>	СН,	4-OC(CH,),	н	143.5—144.5
(113)	CH <sub>3</sub>	СН,	4-SO <sub>3</sub> Na	Н	>350
(114)	O		н	н	260261
(115)	Ô		4-CH,	H	268.5–270
(116)	()		4-OCH <sub>2</sub>	Н	245247
(117)	$\bigcirc$		4-0-(	Н	209211
(118)			4-0(CH <sub>2</sub> ), CH <sub>3</sub>	н	222–225

TABLE (Continued)

	T				
No.	R	R,	х	X,	Melting point uncorrected in °C
(119)	" (*)		4-OC(CH <sub>3</sub> ) <sub>3</sub>	H	203—205
(120)	CH,	CH,	4-80, CH,	н	259-261
(121)	сн,	CH,	4-SO,NH,	н	315317
(122)	CH,	Сн,	3-SO <sub>a</sub> Na	н	> 350
(123)	CH,	Сн,	2-\$0, Na	н	> 350
(124)			4-CH <sub>2</sub>	H	233-236
(125)	$\bigcirc$		4-OCH,	н	236—237.5
126	CH,	Сн,	3-CH <sub>a</sub>	5-CH,	210-212
127	CH <sub>a</sub>	CH,	3-CH,	4-CH,	204—206
128	CH,	CH,	3-OCH <sub>3</sub>	4-0CH,	212-213
129	CH.	CH,	3-0-		223.5–225
130	CH <sub>3</sub>	CH <sub>z</sub>	3-OCH,	4-C,H,	135.5-136.5
131	CH <sub>3</sub>	CH,	3-CH,	4-0CH,	207-208
132	CH <sub>3</sub>	CH,	3-C1	4-C1	272–273
133	CH <sub>3</sub>	CH <sub>3</sub>	2-C1	5-C1	209-211
134	C <sub>2</sub> H <sub>5</sub>	C,H,	4-0Сн,	н	116117
135	C <sub>4</sub> H <sub>9</sub>	C <sub>4</sub> H <sub>9</sub>	4-0CH,	н	10 1-102
136	CH <sub>3</sub>	CH <sub>3</sub>	3-50, Na	4-CH,	>350
137	$\Diamond$		3-SO <sub>s</sub> Na	4-CH,	>350
138	CH <sub>3</sub>	CH3	3-SO <sub>z</sub> Na	4-0CH <sub>3</sub>	>350

	ood (continued)						
No.	R	R <sub>1</sub>	x	X,	Melting point uncorrected in °C		
		<b>~</b> `\					
139		)	3-50, Na	4-OCH,	> 350		
140	Сн.	CH,	3-0C,H,	н	160-161		
141	CH,	СН,	4-0C <sub>2</sub> H <sub>5</sub>	Н	212—213		
142	CH,	Сн,	4-0C2H2OC2H3	н	154—155		
143	CH <sub>3</sub>	сн,	4-OC, H,	н	175—176		
· 144	CH,	сн,	4-00H CH,	Н	162–168		
145	Сн,	СН,	4-C <sub>2</sub> H <sub>6</sub>	н	176—177		
146	CH,	Сн,	4-C₄H,	н	110-112		
147	CH <sub>3</sub>	СН,	4-0C <sub>2</sub> H <sub>4</sub> 0C <sub>3</sub> H <sub>7</sub>	н	118-120		
148	сн,	СН,	3-C <sub>2</sub> H <sub>8</sub>	н	117—118		
149	CH <sub>3</sub>	Сн,	CH,	н	154—156		
150	CH,	CH,	3 0		229-231		
151	сн,	CH3	4-0CH2CH=CH2	H	174175		
152	CH <sub>3</sub>	CH,	4-0CH, CH, OH	н	223-224		
153	CH,	СН,	4-0-i so-C, H,	Ħ	177—178		
154	CH,	CH,	4-0-sec-C <sub>4</sub> H <sub>9</sub>	н	125-127		
155	CH <sub>3</sub>	CH,	4-0C, H, OCH,	н	176-177		
156	сн,	СН,	4-0C,H,OCH(CH,),	н	154155		
157	CH,	СН,	4-0C,H,O-n-C,H,	н	106—107		
158	CH,	СН,	4-OCH CH, OCH,	н	141—142		
159	CH;	СН,	+ос, н, сн осн,	н	143—145		

#### TABLE (Continued)

			T	
R	$\mathbf{R_{i}}$	x	X,	Melting point uncorrected in °C
CH,	CH,	4-OC, H <sub>6</sub> OH	H	
сн,	CH,	4-OC,H,OH	H	
CH,	CH,	4-00H <sub>2</sub> -()	Ħ	179—180
CH,	сн,	4-00 <sub>2</sub> E <sub>4</sub>	Н	,
СН,	CH <sub>3</sub>	3 CH <sub>2</sub> 0 CH <sub>2</sub>		237—239
	CH, CH,	CH, CH, CH, CH, CH, CH,	CH <sub>3</sub> CH <sub>3</sub> 4-OC <sub>3</sub> H <sub>6</sub> OH  CH <sub>3</sub> CH <sub>3</sub> 4-OC <sub>4</sub> H <sub>6</sub> OH  CH <sub>3</sub> CH <sub>3</sub> 4-OCH <sub>2</sub> C  CH <sub>3</sub> CH <sub>3</sub> 4-OCH <sub>2</sub> C  CH <sub>3</sub> CH <sub>3</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>3</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> A-OC <sub>2</sub> H <sub>6</sub> CH  CH <sub>3</sub> CH <sub>2</sub> CH  CH <sub>3</sub> CH  CH  CH <sub>3</sub> CH  CH  CH  CH  CH  CH  CH  CH  CH  CH	CH <sub>3</sub> CH <sub>3</sub> 4-OC <sub>3</sub> H <sub>6</sub> OH H  CH <sub>3</sub> CH <sub>3</sub> 4-OC <sub>4</sub> H <sub>8</sub> OH H  CH <sub>3</sub> CH <sub>3</sub> 4-CCH <sub>2</sub> H

4 - Amino - 2,6 - bis - (piperidino)pyrimidine, which is required for the preparation of compounds (124) and (125), is prepared as follows:

32.8 g of 4 - amino - 2,6 - dichloro-pyrimidine are dissolved in 200 ml of piper-idine, while stirring well, and the temperature rises up to 100° C. After the reaction has subsided, the mixture is refluxed for 12 hours. The mixture containing the piperidine hydro-chloride which has precipitated is poured into water. The oily product which separates out is extracted with methylene chloride, the solution is dried over magnesium sulphate and the methylene chloride is evaporated. The residue is dissolved in other and the solution is saturated with gaseous hydrogen chloride, while stirring well and with cooling. The pale beige hydrochloride which has precipitated is filtered off and dried at 20° C. After recrystallisation from isopropanol with the addition of animal charcoal, 34.4 g (58% of theory) of 4 - amino - 2,6 - bis - (piperidino) - pyrimidine hydrochloride with a melting print of 200 - 200 C. ing point of 269 to 273° C are obtained.

4 - Amino - 2,6 - bis - (diethylamino)-pyrimidine, which is required for the preparation of compound (134) is prepared as

10.0 g of 4 - amino - 2,6 - dichloropyrimidine are heated with 40 ml of diethylamine for 8 hours in an autoclave at 200° C. After cooling to room temperature, the reaction 35 mixture is poured into 200 ml of 10% strength sodium hydroxide solution and the oil which has precipitated is extracted with ether. After

drying and filtering the ether solution, the latter is evaporated and the residue is distilled under a high vacuum and this gives 9.4 g (66% of theory) of 4 - amino - 2,6 - bis-(diethylamino) - pyrimidine with a boiling point of 132 to 137° C/0.07 mm Hg.

If the diethylamine employed in the above example is replaced by di - n - butylamine, this gives 4 - amino - 2,6 - bis - (di - n - butylamino) - pyrimidine, which is required for the preparation of compound (135) and has a boiling point of 160 to 170° C/ 0.05 mm Hg.

1 - Ethoxy - 2 - (4 - aminophenoxy)ethane of the formula

$$\mathbf{H_2N-}\bigcirc -\mathbf{O-}\mathbf{CH_2^-}\mathbf{CH_2^-}\mathbf{OC_2H_5}$$

which is required for the preparation of com-

pound (142), is prepared as follows:
157.6 g of 1-chlor-4-nitrobenzene and
108.1 g of ethylene glycol monochyl ether are dissolved in 300 ml of dimethylsulphoxide, and 123.42 g of a 50% strength aqueous solution of potassium hydroxide are added in the course of 30 minutes, with slight cooling with ice-water. The reaction is weakly exothermic and after the cooling medium has been removed the temperature rises to about 45° C. The mixture is then heated to 65 to 70° C and stirred overnight at this temperature. The reaction mixture is then allowed to run into a mixture of 3 litres of water and ice, while stirring well. The product which has precipitated is stirred ice-cold for a further 15

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minutes and is then filtered off, washed with water and dried in vacuo. The yield of 1ethoxy - 2 - (4 - nitrophenoxy) - ethane is 179 g (85% of theory). Melting point = 67 to 68° C.

1 - Ethoxy - 2 - (4 - nitrophenoxy) - ethane is reduced to 1 - ethoxy - 2 - (4 - aminophenoxy) - ethane with Raney nickel by the conventional route.

The amine is a pale yellow oil. Boiling point u.os tmm 110° C.

Example 2.

1 g of the brightener of the formula (100) is dissolved in 1,000 ml of dimethylformamide. 100 ml of water which contain 0.06 g of an alkyl polyglycol ether are added to 3 ml of this solution. Polyamide fabric (polyamide 6 or 66) weighing 3 g is added to this brightener solution, which has been warmed to 60° C. The temperature is raised to 95 to 97° C in the course of 10 to 15 minutes and this temperature is maintained for 30 minutes. The fabric is then rinsed in running cold water for 2 minutes and then dried for 20 minutes at 60° C.

The fabric treated in this way shows a strong brightening effect with good farmers to light.

Example 3.

1 g of the brightener of the formula (100) is 30 dissolved in 1,000 ml of dimethylformamide. 100 ml of water which contain 0.12 ml of 85% strength formic acid are added to 3 ml of this solution. Polyamide fabric (polyamide 6 or 66) weighing 3 g is added to this brightener solution, which has been warmed to 60°C. The temperature is raised to 95 to 97° C in the course of 10 to 15 minutes and this temperature is maintained for 30 minutes. The fabric is then rinsed in running cold water for 2 minutes and subsequently is dried for 20 minutes at 60° C.

The fabric treated in this way shows a strong brightening effect with good fastness to light.

Example 4.

Using a liquor ratio of 1:20, polyamide fibre fabric (for example "Perlon-Helanca":
Registered Trade Mark), is washed for 15
minutes in a liquor, at 55° C, which contains, per litre, 0.002 g of a brightener of the formula (100) and 4 g of a washing agent of the following composition: 15.7% of an alkyl aryl-sulphonane, 3.7% of a fatty alcohol sulphate, 2.7% of coconut acid monoethanolamide, 39.0% of sodium tripolyphosphate, 4.0% of sodium silicate, 2.0% of magnesium silicate, 1.0% of carboxymethylcellulose, 0.5% of the sodium salt of ethylenediaminetetraacetic acid (EDTA) and 6.7% of water, made up to 100% with sodium sulphate.

The fabric is then washed under running water for & minute and dried for 20 minutes at 60° C in a drying cabinet.

The fabric shows a strong brightening effect with good fastness to light.

The washing agent of the above composition can also contain the brightener of the formula (100) as a directly incorporated substance.

Example 5. 1 g of the brightener of the formula (100) is dissolved in 1,000 ml of dimethylformamide. 95 ml of water which contain 0.06 ml of 40% strength acetic acid are added to 6 ml of this solution. Cellulose acetate fabric weighing 3 g is added to this brightener solution, which has been warmed to 40° C. The temperature is raised to 75 to 80° C in the course of 10 to 15 minutes and this temperature is maintained for 30 minutes. The fabric is then rinsed in running cold water for 2 minutes and subsequently is dried at 60° C for 20 minutes.

The fabric obtained in this way shows a 85 strong brightening effect with good fastness to

Example 6. 1 g of the brightener of the formula (100) is dissolved in 1,000 ml of dimethylformamide. 95 ml of water which contain 0.06 ml of 40% strength acetic acid are added to 6 ml of this solution. Cellulose triacetate fabric weighing 3 g is added to this brightener solution, which has been wanned to 40° C. The temperature is raised to 95 to 97° C in the course of 10 to 15 minutes and this temperature is maintained for 30 minutes. The fabric is then rinsed in running cold water for 2 minutes and subsequently is dried for 20 100

minutes at 60° C. The fabric obtained in this way shows a strong brightening effect with good fasmess

to light.

Example 7. 105 In a kneader, 67 parts of polyvinyl chloride powder, 33 parts of dioctyl phthalate, 2 parts of di-n-butyl dilauryl-dioxystannate and 0.3 part of sodium pentaoctyl-tripolyphosphate are gelatinised with 0.05 part of the optical 110 brightener of the formula (101) on mixing rolls at 160° C for 15 minutes and the mixture is then drawn out to films. The polyvinyl chloride film produced in this way has a strong fluorescence and a brilliant white appearance 115 in daylight.

Example 8. 1,000 parts of polyamide chips obtained in a known manner from hexamethylene adipate are mixed in a drum with 5 parts of titanium dioxide and 0.5 part of the optical brightener

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of the formula (101) for 10 to 16 hours. The chips treated in this way are melted in an apparatus, with the exclusion of oxygen, and the melt is stirred briefly. The melt is then spun out, under a nitrogen atmosphere of 5 atmospheres gauge, through spinnerets and stretched. The polyamide filaments obtained in this way have a high whiteness.

WHAT WE CLAIM IS:-1. A v - triazolyi[4,5 - d] - pyrimidine 10 of the formula

in which Q and Q1 independently of one another are a secondary or tertiary amino radical; X is hydrogen, alkyl having 1 to 6 carbonatoms, alkoxy having 1 to 6 carbon atoms, allyloxy, hydroxyalkoxy having 2 to 4 carbon atoms, alkoxyalkoxy having a total of 3 to 8 carbon atoms, benzyloxy, phenethyloxy, halogen, phenoxy, phenoxyalkoxy having 1 to 3 carbon atoms in the alkoxy part, sulpho, cyano, —COOY (in which Y is hydrogen, a salt-forming cation or alkyl having 1 to 18 carbon atoms) or

## $-SO_2NY_1Y_2$ or $-CONY_1Y_2$

(in which Y<sub>1</sub> and Y<sub>2</sub> independently of one another are hydrogen or alkyl having 1 to 4 carbon atoms); X1 is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 arbon atoms, halogen, sulpho, cyano, -COOY (in which Y is hydrogen, a saltforming cation or alkyl having 1 to 18 carbon atoms) or

#### -SO,NY,Y, or -CONY,Y,

(in which Y1 and Y2 independently of one another are hydrogen or alkyl having 1 to 4 carbon atoms), or X and X, together form a methylenedioxy, ethylenedioxy or methyleneoxymethyleneoxy radical; and X, is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms or halogen

2. A v - triazolyl [4,5 - d] - pyrimidine according to claim 1, of the formula

(2) 
$$R_{2} = R_{1}$$

$$R_{2} = R_{1}$$

$$R_{3}$$

in which R and R2 independently of one another are hydrogen, alkyl having 1 to 12 carbon atoms, hydroxyalkyl having 2 to 6 carbon atoms or dialkylaminoalkyl having 2 to 4 carbon atoms per alkyl part; R, and R, independently of one another are alkyl having 1 to 12 carbon atoms or hydroxyalkyl having 2 to 6 carbon atoms; or R<sub>1</sub> together with R, or R<sub>2</sub> together with R<sub>2</sub>, form with the nitrogen atom to which they are attached a 5-membered or 6-membered saturated heterocyclic structure which can contain one or more further hetero-atoms; X is hydrogen, alkyi having 1 to 6 carbon atoms, alkoxy having 1 to 6 carbon atoms, allyloxy, hydroxy-alkoxy having 2 to 4 carbon atoms, alkoxyalkoxy having a total of 3 to 8 carbon atoms, benzyloxy, phenethoxy, halogen, phenoxy, phenoxyalkoxy having 1 to 3 carbon atoms in the alkoxy part, sulpho, cyano, —COOY, or —SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or —CONY<sub>1</sub>Y<sub>2</sub>; and X<sub>1</sub> is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms, halogen, sulpho, cyano, —COOY, or X and X<sub>1</sub> together form a methylenedioxy, ethylenedioxy or methyleneoxymethyleneoxy radical.

3. A v - triazolyl [4,5 - d] pyrimidine

according to claim 2, of the formula

$$\begin{array}{c}
R_1^1 \\
R_2^1
\end{array}$$

$$\begin{array}{c}
R_1^1 \\
R_3^1
\end{array}$$

$$\begin{array}{c}
R_1^1 \\
R_2^1
\end{array}$$

in which R' and R2' independently of one another are hydrogen, alkyl having 1 to 4 carbon atoms or hydroxyalkyl having 1 to 4 carbon atoms; R<sub>1</sub>' and R<sub>8</sub>' independently of one another are alkyl having 1 to 4 carbon atoms or hydroxyalkyl having 2 to 6 carbon atoms; or R<sub>1</sub>' together with R', or R<sub>2</sub>' together with the nitrogen atom to which they are attached from a atom to which they are attached, form a 5-membered or 6-membered saturated heterocyclic structure which can contain a further nitrogen atom or an oxygen atom; X' is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms, allyloxy, hydroxyalkoxy having 2 to 4 carbon atoms, alkoxyalkoxy having a total of 3 to 8 carbon atoms, benzyloxy, phenethoxy, chlorine, phenoxy, phenoxypropoxy, sulpho, cyano, —COOY, —SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or —CONY<sub>1</sub>Y<sub>2</sub>; and X<sub>1</sub>" is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms or chlorine, or X1" and X' together form a methylenedioxy, ethylenedioxy or methyleneoxymethyleneoxy radical.

4. A v - triazolyl[4,5 - d]pyrimidine according to claim 3, of the formula

in which R" and R<sub>2</sub>" independently of one another are hydrogen, alkyl having 1 to 4 carbon atoms or hydroxyethyl; R," and R," independently of one another are alkyl having I to 4 carbon atoms or hydroxyethyl; or R,'' together with R'', or  $R_n''$  together with  $R_2''$ , and the nitrogen atom to which they are attached are pyrrolidino, piperidino, morpholino, N'-hydroxyethylpiperazino or N'-hydroxyethylpiperazino quaternised by alkyl having I to 4 carbon atoms; and X" is hydrogen, methyl, chlorine, alkoxy having 1 to 4 carbon atoms, allyloxy, hydroxyalkoxy having 2 to 4 carbon atoms, alkoxyalkoxy having 3 to 6 carbon atoms, benzyloxy, phenoxy, phenoxy-propoxy, sulpho, cyano, —COOY (in which Y is hydrogen, a salt-forming cation or alkyl having 1 to 4 carbon atoms), —SO<sub>3</sub>NY<sub>1</sub>Y<sub>2</sub>, or —CONY<sub>1</sub>Y<sub>2</sub>; X<sub>1</sub>"' is hydrogen, alkyl hav-20 ing 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms or chlorine, or X" and X," together form a 3,4-methylenedioxy, 3,4ethylenedioxy or 3,4-methyleneoxymethylene-25 oxy radical.

5. A v - triazolyl [4,5 - d] pyrimidine according to claim 4, of the formula

in which R" is alkyl having 1 to 4 carbon atoms, or two adjacent R", together with the nitrogen atom to which they are attached, are piperidino or morpholine; X" is hydrogen, methyl, allyloxy, hydroxyethoxy, alkoxyalkoxy having a total of 3 to 6 carbon atoms, benzyloxy, alkoxy having 1 to 4 carbon atoms, phenoxy, alkoxy having 1 to 4 carbon atoms, phenoxy, phenoxypropoxy or sulpho, and X<sub>1</sub>"" is hydrogen, alkyl having 1 to 4 carbon atoms, phenoxy, phenoxypropoxy or sulpho, and X<sub>1</sub>"" is hydrogen, alkyl having 1 to 4 carbon atoms or methoxy, or X<sub>1</sub>"" and X" together is 3,4-methylenedioxy, 3,4-ethylenedioxy or 3,4-methyleneoxymethyleneoxy.

6. Each of the compounds (100) to (125) hereinbefore identified.

45 7. Each of the compounds (126) to (164) hereinbefore identified.

8. A process for the preparation of a v-triazolyl[4,5-d] pyrimidine as defined in claim 1 which comprises coupling an amine of the formula

in which Q and Q<sub>1</sub> are as defined in claim 1, with a diazenium salt of the formula

in which Z is an anion and X to X, are as defined in claim I, and oxidatively cyclising the resulting o-aminoazo compound of the formula

in which Q, Q<sub>1</sub> and X to X<sub>2</sub> are as defined in claim 1, at a temeprature of from 70° to 100°C.

9. A v-triazoly1[4,5-d]pyrimidine according to claim 1 when prepared by a process claimed in claim 8.

10. A process for the optical brightening of organic material, which comprises applying to or incorporating in the material a v-triazolyl[4,5-d]pyrimidine as claimed in any one of claims 1 to 5, 7 or 9.

11. A process according to claim 10 wherein the material is polyamide, cellulose acetate, cellulose triacetate or polyvinyl chloride material.

12. A process according to claim 10 or 11, wherein 0.001 to 2% by weight of the brightener, relative to the weight of the material to be brightened, is used.

13. Organic material optically brightened by a process claimed in claim 10 or 11.

14. Organic material containing 0.001 to 2% by weight of a v-triazolyl[4,5-d]-pyrimidine as claimed in any one of claims 1 to 5, 7 or 9

1 to 5, 7 or 9.

15. The use of a v-triazolyl[4,5-d]pyrimidine as claimed in any one of claims
1 to 5, 7 or 9 as an optical brightener for
organic material.

16. A v-triazolyl[4,5-d] pyrimidine according to claim 1, wherein X is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms, halogen, phenoxy, phenoxy-alkoxy having I to 3 carbon atoms in the alkoxy part, sulpho, cyano, —COOY, —SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or —COY<sub>1</sub>Y<sub>2</sub> or X and X<sub>1</sub> together form a methylenedioxy or ethylenedioxy radical.

17. A v-triazolyl[4,5-d]pyrimidine according to claim 3, wherein X' is hydrogen, alkyl having 1 to 4 carbon atoms, alkoxy having 1 to 4 carbon atoms, chlorine, phenoxy, phenoxypropoxy, sulphn, cyano, —COOY, —SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or —CONY<sub>1</sub>Y<sub>2</sub> or X' and X<sub>1</sub>" together form a methylenedioxy or ethylenedioxy radical.

18. A v-triazoly[4,5-d] pyrimidine according to claim 4, wherein X" is hydrogen, methyl, chlorine, alkoxy having 1 to 4 carbon atoms, phenoxy, phenoxypropoxy, sulpho, cyano, —COOY, —SO<sub>2</sub>NY<sub>1</sub>Y<sub>2</sub> or —CONY<sub>1</sub>Y<sub>2</sub>, X<sub>1</sub>" is hydrogen, methyl, 15 alkoxy having 1 to 4 carbon atoms or chlorine, or X" and X<sub>1</sub>" together form a 3,4-methylenedioxy or 3,4-ethylenedioxy radical.

A process for the optical brightening of organic material, which comprises applying to or incorporating in the material a compound claimed in claim 6, 16, 17 or 18.

20. A process according to claim 19 wherein the material is a polyamide, cellulose ester or polyvinyl chloride.

21. A process according to claim 19 or 20, wherein 0.001 to 2% by weight of the brightener, relative to the weight of the brightener, is used.

22. Organic material optically brightened by a process claimed in claim 19, 20 or 21.

23. The use of a v-triazolyl[4,5-d]-pyrimidine claimed in claim 6, 16, 17 or 18 as an optical brightener for organic material.

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